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LTD

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(54) LITHIUM BATTERY

(57) Abstract:

PURPOSE: To stabilize voltage and current and improve reliability by forming an electrolyte from a specified high molecular copolymer cross-linked body in which an electrolyte is connected to a silicon atom through a hydrocarbon group. CONSTITUTION: In a lithium battery in which negative and positive electrodes having lithium metal, a lithium alloy or a lithium inserting type carbon as active materials are constituted through an electrolyte, the electrolyte is a high molecular copolymer cross-linked material consisting of a high molecular compound containing a polyorganosiloxane chain containing a lithium carboxylate group represented by the formula I in which the electrolyte is connected to a silicon atom through a hydrocarbon group and a polyoxyalkylene chain. The kind, form, and crosslinking density of the high molecular compound are not particularly limited.



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(54)【発明の名称】 リチウム電池

(57)【要約】

【目的】 安定した電圧、電流を与える全固体リチウム 電池を提供する。

【構成】 金属リチウム. リチウム合金もしくはリチウム挿入型炭素を活物質とする負極と. 正極とが電解質を介して構成されているリチウム電池において、前記電解質が. 炭化水素基を介してケイ素原子に結合した式、一〇〇〇 Li' で示されるリチウムカルボキシレート基を含有するボリオルガノシロキサン鎖とボリオキシアルキレン鎖含有高分子化合物とから構成された高分子共量合体架橋物からなることを特徴とするリチウム電池。

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【特許請求の範囲】

【論求項1】 金属リチウム、リチウム合金もしくはリチウム挿入型炭素を活物質とする負極と、正極とが電解質を介して構成されているリチウム電池において、前記電解質が、炭化水素基を介してケイ素原子に結合した式、一COO・L・・で示されるリチウムカルボキシレート基を含有するポリオルガノシロキサン鎖とポリオキシアルキレン鎖含有高分子化合物とから構成された高分子共重合体架橋物からなることを特徴とするリチウム電池。

【請求項2】 高分子共重合体架績物が、(A)1分子中に少なくとも2個のカルボキシル基含有炭化水素基を有するポリオルガノシロキサン、(B)1分子中に少なくとも2個の水酸基を有するポリオキシアルキレン鎖含有高分子化合物、および(C)リチウム金属原子を含むアルカリ化合物からなる混合物を脱水縮台反応により硬化せしめることにより製造されたものである請求項1記載のリチウム電池。

【発明の詳細な説明】

[0001]

【産業上の利用分野】本発明はリチウム電池、詳しくは 全固体で安定した電圧、電流を与えるリチウム電池に関 する。

[0002]

【従来の技術および発明が解決しようとする課題】近 年、リチウム電池はマイクロエレクトロニクスの発展に 住い、高性能化、小形化、薄型化が一段と進んでいる。 かかるリチウム電池においては負極、正極の材料および 形態の改良もさることながら、それに組み込まれる電解 質の改良が要求されており、とりわけ固体化、高柔軟 性、高成形加工性を与える意味で固体電解質が重要にな ってきている。さらに、高イオン伝導性、高リチウムイ オン輸率、高信頼性、耐湿性等種々の高度特性が要求さ れている。従来、このような電解質としては、高分子樹 脂マトリックスにリチウム塩を溶解、分散させた固体電 解質材料等が知られている。中でもポリオルガノシロキ サン鏡とポリエチレンオキシド (PEO) 鎖とを組み合 わせた固体電解質の台成が盛んに試みられている。例え は、特開昭62-209169号公報にはシロキサンと PEOとを白金触媒によるヒドロシリル化反応や放射線 (電子線等) 照射により架橋させ、得られた架橋物にリ チウムイオンを分散させて、固体電解質としたものが開 示されている。また、特別昭63-170857号公報 にはこれらの固体電解質を組み込んだリチウム電池が開 示されている。しかし、これらの方法では、2種以上の 原料を相溶させるために有機溶剤を用いる必要があっ た。有機溶剤の使用は一部の製品には適用できないばか りか工程の煩雑化を招き、また作業環境の悪化、周辺材 料の損傷、最終生成物への有機溶剤の残留等を起こす恐 れがあった。また有機溶剤により原料が相溶できても、

有機溶剤が蒸発する過程などで相分離を起こす等により 架橋反応の完結性が完全には保証されず、結局製品の品 質の低下、再現性のなさ等に結び付くため実用化には問 題があった。さらに、特別平2-230667号公報に はポリエチレンオキシド鎖をグラフトしたポリスチレン にリチウム塩を分散させた固体電解質を組み込んだリチ ウム電池が開示されている。この方法においては材料の 均質性は向上するものの、以下の欠点があった。すなわ ち、特開昭63-170857号公報、特開平2-23 10 0667号公報等で述べられているリチウム塩を分散さ せる形式の固体電解質においては、電気伝導によりリチ ウムイオンと同時に負イオンの移動が起こるものであっ た。負イオンの移動は多くの場合弊害となることが多 い。例えば、充放電時に負イオンの移動により電解質内 に分極が起こることにより電極/電解貿界面の構造が経 時的に変化し、金属イオンの移動に伴う定常電流が流れ なくなる。負イオンは正イオンとの電荷補償のために存 在せざるを得ないが、その移動度をなるべく低下させる こと、すなわち、正イオン単独伝導型固体電解質の作製 20 が試みられている。例えば、日本化学会1988春期年 会講演費号2XIIC08には、オクタメチルシクロテト ラシロキサンとベンゼンスルホン酸メチルエステルをプ ラズマ重合させた後、PEOを複合させ、さらにヨウ化 リチウムで処理することによりスルホネートイオンが固 定されたリチウムイオン単独伝導型固体電解質薄膜が開 示されている。しかし、この場合、プラズマ重合は、反 応が複雑なため意図する高分子の構造を得ることは困難 であり、完全なリチウムイオン単独伝導型固体電解質に はならないという欠点があった。さらに、プラズマ宣合 30 のため、応用範囲に制約があるという欠点があった。こ のように、現在まで提案されたリチウム電池は、たとえ 固体電解質を組み込んだ場合であっても、いずれも固体 電解貿自体の特性あるいは製造方法に欠点を有していた ため、電池の特性および製造法が満足すべきものではな

【0003】本発明者らはかかる問題点を解決すべく観意検討した結果、特定の高分子共宣合体架橋物中にリチウムイオンを分散させてなる材料が、上記のような欠点がなくイオン伝導性に優れ、しかも組成を操作することにより正イオン単独伝導型固体電解質材料になり得、リチウム電池の電解質として組み込んだ場合、上記のような欠点を払拭できることを見出し本発明に到達した。本発明の目的は安定した電圧、電流を与える全固体リチウム電池を提供するにある。

[0004]

【課題の解決手段とその作用】かかる本発明は、金属リチウム、リチウム合金もしくはリチウム挿入型炭素を活物質とする負極と、正極とが電解質を介して構成されているリチウム電池において、前記電解質が、炭化水素基50を介してケイ素原子に結合した式、-COO'L' で

示されるリチウムカルボキシレート基を含有するポリオルガノシロキサン鎖とボリオキシアルキレン鎖含有高分子化合物とから構成された高分子共重合体架振物からなることを特徴とするリチウム電池に関する。

【0005】とれについて説明するに、本発明でいうり チウム電池とは全固体である一次電池および充放電可能 な二次電池双方を指す。一次あるいは二次を決定する要 因としては主に正極活物質の種類である。本発明では、 リチウム電池の高性能化の1つとして先に述べたリチウ ムイオン単独伝導性を目的としているため、リチウム電 10 他の形式としては、一次、二次どちらにおいてもいわゆ るロッキングチェア型の帶成にする必要がある。すなわ ち、正負両極における電池反応を司どるのはリチウムイ オンのみとするものである。従って正極においては、リ チウムイオンが挿入(放電時)、脱離(二次電池におけ る充電時) する形式である必要がある。電池の常造とし ては負極-固体電解質-正極のサンドイッチ構造を有す るものであり、大きさ、厚さおよび形態は限定しない。 が、典型的には面積数ork、厚さ約1mm程度のペーパー 型からコイン型、スパイラル方式のシリンダー型まで多 20 様のものが可能である。負極は代表的にはリチウム金属 箔を用いるが、負極活物質としてのリチウム金属が電極 反応に有効な形態で存在している導電性シートであれば 特に限定されない。例えば、カーボンシートにリチウム などが保持されたもの、あるいはリチウムーアルミニウ ム合金などが挙げられる。正極の活物質としては、上記 電池反応を起こし得るものであれば特に限定されない が、典型的には無機層状化合物を用いる。例えば、二酸 化マンガン、酸化ヴァナジウム、二硫化チタン、酸化コ バルト、酸化ニッケル、硫化モリブデンやこれらの複合 30 体等を用いる。これらの化合物は通常、脆く、また導電 性が低いために、有機樹脂を粘結材として用い板状にす る工夫がされたり、導電性付与剤としてカーボン粒子等 が添加される。また、特に二次電池を構築する場合は、 充放電のサイクルによる層状構造の不変性あるいはリチ ウムイオンの吸脱者性等から一部リチウム原子を構造に 取り込ませた層状化合物を用いると好道である。

【0006】次いで上記高分子共宣合体架橋物について 説明するに、この架橋物はメチレン基、エチレン基、プ ロビレン基、プチレン基、ペンチレン基、ヘキセレン 基、ヘプチレン基。オクチレン基等の炭素数1~8のア ルキレン基;フェニレン基、ナフチレン基等の炭素数8 ~20のアリーレン基で例示されるような炭化水素基を 介してケイ素原子に結合した、式、-COO・L 1・で示 される化学構造を有するリチウムカルボキシレート基合 有ポリオルガノシロキサン鎖とポリオキシアルキレン鎖 含有高分子化合物の種類、形態、架橋密度等については 特に限定されない。

【0007】本発明の固体電解質は前述のような負イオ 50 両末端がトリメチルシロキシ基で封鎖されたメチルカル

ンを含む基 (~COO*) が炭化水素基を介してケイ素原子に結合されているため、負イオンの移動度は極端に低い。従って、この高分子共量合体架橋物中ではリチウムイオンが負イオンから解離し、リチウムイオンが移動することによりイオン導電性が発現する。高分子共量合体架橋物内に存在する負イオンの種類が一COO*のみであれば、本固体電解質材料はリチウムイオン単独伝導体として働く。

【0008】かかる固体電解質の製造方法について好ましい方法は、以下の方法が推奨される。「(A)1分子中に少なくとも2個のカルボキシル基含有炭化水素基を有するポリオルガノシロキサン、(B)1分子中に少なくとも2個の水酸基を有するポリオキシアルキレン鎖含有高分子化合物、および(C)リチウム金属原子を含むアルカリ化合物からなる混合物を脱水縮合反応により硬化せしめることにより製造する方法。」

【0009】 これについて説明するに、(A)成分のポリ オルガノシロキサンは、高分子共量合体架積物を形成す るためには1分子中に2個以上のカルボキシル基含有炭 化水素基を有することが必要であり、また、このカルボ キシル基含有炭化水素基が結合したシロキサン単位とそ れ以外のシロキサン単位の比率が 0.01~100の範 囲内にあるものが好ましい。本成分の分子構造は直鎖 状、分岐状、環状、網状、三次元構造の何れでもよい が、高分子共重合体架積物の形成の容易さからその半数 以上は直鎖状もしくは分岐状であることが好ましい。ま た。その分子量は特に限定されないが、製造の容易さ、 高分子共宣合体架積物としての適度の硬さを得るために は100~100万の範囲内にあることが好ましい。 (A)成分のカルボキシル基含有炭化水素基としては、例 えば、一般式HOOC-R¹-(式中、R¹はメチレン 基、エチレン基、プロピレン基、プチレン基、ペンチレー ン益、ヘキシレン基、ヘプチレン基、オクチレン基等の 炭素数1~8のアルキレン益またはフェニレン益、ナフ チレン基等の炭素数6~20のアリーレン基である。) で示される基が挙げられ、とれらの中でもカルボキシア ルキル基が好ましく、カルボキシプロピル基が特に好ま しい。(A)成分中のカルボキシル基含有炭化水素基以外 の有機基としては、例えば、メチル基、エチル基、プロ 40 ピル基等のアルキル基:フェニル基、トリル基、キシリ ル益等のアリール基:ベンジル基,フェネチル益等のア ラルキル基が例示される。また、ケイ素原子に結合した 基としては、少量の水素原子、アルコキシ基が含まれて もよい。経済性および良好な高分子共重合体架橋物の形 成性の観点からはケイ素原子に結合したオルガノ基の半 数以上はメチル益であることが好ましい。かかるポリオ ルガノシロキサンとしては、例えば、分子鎖両末端がト リメチルシロキシ基で封鎖されたメチルカルボキシブロ ビルシロキサン・ジメチルシロキサン共宣合体、分子鎖

ボキシプロピルシロキサン・メチルフェニルシロキサン 共重合体が挙げられる。 かかるポリオルガノシロキサン の合成方法としては程々の方法が知られているが、その 1つの方法としてはシアノ基を有するオルガノジクロロ シランとシアノ差を有さないオルガノジクロロシランを 共加水分解して得られる環状物と末端停止剤とを確酸水 溶液中で撹拌し、シアノ益がカルボキシル基に転化する 反応と開環宣合を起こさせる方法が挙げられる。

【0010】(B)成分の高分子化合物は上記(A)成分の 架橋剤であり、架橋剤としての働きをするためには、1 分子中に少なくとも2個の水酸基を含有することが必要 である。また、高イオン伝導性発現のため高分子共量合 体架橋物中にはポリオキシアルキレン鎖を有することが米 *好ましいが、そのためには(B)成分の高分子化合物構造 中にポリオキシアルキレン鎖を含有することが必要であ る.

【0011】ところで、(A)成分と(B)成分が互いに相 溶することは工程中に溶剤を使用することが避けられる ばかりか、架橋反応の完結、架橋生成物の構造、物性の 良好な再現性をもたらすため大変重要なことである。 (A)成分と相溶するための(B)成分の分子構造および化 学常造についてはシロキサン単位を有していることが好 ましく、高分子共宜合体架橋物の形成の容易さ等から一 砂式

[12]

(式中、R'は1価の有機基、R'は2価の有機基、R1 はアルキレン芸、R'は1価の有機基、a、cは0~1 000の整数、bは2~1000の整数, pは2~10 0の整数である。) で示される末端に水酸基を有するポ リオキシアルキレン鎖を少なくとも2本グラフト鎖とし て有するポリオルガノシロキサンが好ましい。 かかるオ ルガノポリシロキサンは上式中、R1は、メチル基、エ チル基、プロビル基等のアルキル基;フェニル基、トリ ル基、キシリル基等のアリール基;ベンジル基、フェネ チル基等のアラルキル基が例示される。また、一部、少 および良好な高分子共重合体架橋物の形成性の観点から はR'の半数以上はメチル基であることが好ましい。R' はメチレン基、エチレン基、プロピレン基、ブチレン 基、ベンチレン益、ヘキシレン基、ヘブチレン益、オク チレン基等の炭素数1~8のアルキレン基またはフェニ レン益、ナフチレン基等のアリーレン基等の2価の有极 基である。R*はメチレン基、エチレン基、プロピレン 基、ブチレン基、ペンチレン基、ヘキシレン基、ヘプチ レン益等のアルキレン基である。R'はメチル基、エチ ル基、プロビル基等のアルキル基:アセチル基またはプ 40 ロビオニル基等のアシル基である。a. cは0~100 ○の範囲内であり、bは2~1000の範囲内であり、 それぞれ特に限定されないが、(A)成分と(B)成分が相 溶するためにはaと(b+c)の比率は(1:5)~ (5:1)の範囲内が好ましい。

【0012】かかるグラフト共宣合体の合成方法として は種々の方法が知られているが、その1つの方法として は側鎖の一部が水素原子で置き換わったポリオルガノシ ロキサンに、片末端に不飽和炭化水素基を有し他末端に アシロキシ基を有するポリオキシアルキレンと、片末蟾 50 る。しかし、(A)成分と(B)成分が钼溶する条件にお

20 に不飽和炭化水素基を有し他末端にトリメチルシリル基 を有するポリオキシアルキレンを所定の比率によりヒド ロシリル化反応によりグラフトさせ、しかる後に過剰量 のアルコールにより グラフト末端のトリメチルシリル基 のみを水酸基に転化する方法が挙げられる。

【0013】(B)成分は上記のような1分子中に少なく とも2個の水酸基を有し、 構造中にポリオキシアルキレ ン鎖を含む高分子化合物であるが、より高いイオン伝導 性を要求される場合には、この高分子化合物は一般式目 O- (R°O) q-H (式中、R*はアルキレン益、qは 量の水素原子、アルコキン基が含まれてもよい。経済性 30 1~100の整数である。)で示される分子鎖両末端に 水酸基を有するポリオキシアルキレンまたはHO-(R 'O) r-R'(式中、R'はアルキレン基、R'は1価の 有機基, r は 1 ~ 1 0 0 の整数である。)で示される分 子鎖片末端に水酸基を有するポリオキシアルキレンであ ることが好ましい。かかるポリオキシアルキレンの上式 のR¹, R⁷はメチレン基、エチレン基、プロピレン基。 ブチレン基、ペンチレン基、ヘキシレン基、ヘブチレン 基等のアルキレン基である。R'はメチル基、エチル 基、プロビル基等のアルキル基:アセチル基またはプロ ピオニル基等のアシル基である。q、rは1~100の 範囲内であり、5~20の範囲内が好ましい。かかる分 子鎖両末端に水酸基を有するポリオキシアルキレンまた は分子鎖片末端に水酸基を有するポリオキシアルキレン は上記(A)成分と縮合反応し、高分子共全合体架橋物中 にポリオキシアルキレンの架橋鎖またはグラフト鎖の含 有量を増加する働きをする。本発明においては、かかる 架橋鎖またはグラフト鎖が存在しなくても高分子化合物 中のポリオキシアルキレン鎖によりポリオキシアルキレ ンの架積鎖またはグラフト鎖を形成させることはでき

いては高分子化合物中のポリオキシアルキレン鎖の導入 量には限界があるため、分子鎖両末端に水酸基を有する ボリオキシアルキレンまたは分子鎖片末端に水酸益を有 するポリオキシアルキレンを(B)成分中に含有させる方 が好ましい。また本発明においては、高分子共重合体架 **橋物中にポリオキシアルキレンのグラフト鎖が若干存在** した方がイオン伝導性が向上する傾向にある。

【0014】(C)成分は、リチウム原子を含むアルカリ 化合物であり、その積額は特に限定されないが、水酸化 物、アルコラート、水素化物等が好ましく、特に水酸化 10 物の形態、すなわち水酸化リチウム (LiOH) が使用 されることが好ましい。かかる(C)成分は、(A)成分と (B)成分のエステル化反応の触媒作用をすると同時に、 最終的には(A)成分中のカルボキシル基を脱水によりリ チウムカルボキシレート化する役割をする。その結果、 高分子共宣合体架織物としては、負イオン(カルボキシ レートイオン) はシロキサンポリマー鎖に固定され、対 イオンである正イオン(リチウムイオン)が分散された 形態になる。また、その分散量は高分子共重合体架橋物 ウムイオンのモル数:[Li*]の比率比[Li*]/[R 〇] が0.005~0.25とすることが好ましく。より 好ましくは0.02~0.1である。これはこの比が 0. 25を越えると高分子共重合体架橋物の極性が上がり、 セグメントの運動性が悪くなり、また 0.005 未満に なるとキャリア数の低下から高いイオン伝導性が得難く なるからである。また、(A)成分中のカルボキシル基の モル敷と、(B)成分中の水酸基のモル数と(C)成分中の アルカリ当量の合計の比率は(1:10)~(10: 0) の範囲内が好ましい。 これは (A) ~(C)成分はい ずれの比率でもエステル化反応が起こり、固体化した高 分子共宜台体架橋物が得られるが、(B)成分中の水酸基 のモル数と(C)成分中のアルカリ当量の合計と(A)成分 中のカルボキシル基のモル数の比が大幅に異なると、固 体電解質材料中に未反応のカルボキシル基または水酸基 が残存した状態となる傾向にある。この状態での固体電 解買を電池に組み込んだ場合には、電極材との反応等の 弊害が起こることが予想されるので、好ましくない。ま た. 上記のモル比が1になれば、正イオン程はリチウム 40 イオンのみ、負イオン種は固定されたカルボキシレート イオンのみとなり、完全なリチウムイオン単独伝導型固 体電解質となる。

【0015】この方法においては(A)成分~(C)成分か らなる混合物を脱水縮合反応により硬化させ高分子共重 台体架橋物を形成させるのであるが、この架橋物は主に (A)成分中のカルボキシル昔と(B)成分の水酸益とのエ ステル化反応によって形成されるものである。その反応 手段としてはカルボキシル基と水酸基との反応に用いら

ステル化反応触媒を使用し、常温または加熱下に反応さ せる方法が利用できる。また、加熱はこのエステル化反 応を促進すると同時にエステル化反応によって発生した 副生物である水を除去するために有効な技術手段であ り、その温度は通常150℃以下である。また、この架 錯反応は無溶媒の条件で行うことができる。(A)成分と (B)成分は攪拌等の操作で容易に均一に混合できる。 (C)成分はオキシアルキレン鎖に対して溶解性を持つの で、(B)成分に予め溶解させておくか、(A)成分と(B) 成分を混合させた後に添加してもよい。(C)成分のオキ シアルキレン鎖への溶解は撹拌等の操作でも可能だが、 溶解時間短縮のためには加熱あるいは超音波照射等の操 作あるいはごく少量の水の添加が効果的である。また、 溶解工程において有機溶剤の使用が許容できる場合に は、有機溶剤中で(A)成分~(C)成分を混合、溶解さ せ、しかる後に溶剤を蒸発させてもよい。かかる有機溶 剤は特に限定されないが、例えば、テトラヒドロフラ ン、ジオキサン、アセトニトリル、ジメチホルムアミ ド、ジメチルスルホキシドが挙げられる。また、このエ 中のオキシアルキレン基のモル数 [RO] に対するリチ 20 ステル化反応は副生物として水を生成するので、最終的 には(A)成分~(C)成分の混合体を減圧下に置くことが 好ましく、例えば、常圧加熱下でエステル化反応をある 程度進行させた後に減圧下で加熱し、生成した水の除去 とエステル化反応の完結を同時に行う方法が推奨され る。ただし、溶解工程に溶剤を使用した場合は、常圧下 に有機溶剤の沸点以下でエステル化反応をある程度進行 させた後に、有機溶剤を蒸発させ、しかる後に減圧下で 加熱する必要がある。

【0016】本発明のリチウム電池は上記の正極、負 1) の範囲内であり、(1.0:1,2)~(1.2:1. 30 極. 固体電解質を組み合わせてなるものであるが. その 構築法については特に限定しない。 各々を別々に作製し てから3者を組み合わせてもよい。しかし、一般的に界 面における接触面積が大きいほど電池としての効率が高 まることから、正極上に固体電解質原料の混合物をキャ ストし、架橋反応を行わせてフィルム化した後、負極を 張り合わせる方法が推奨される。また、水分により、負 極、固体電解質は変質を受けるため、電池の作製は乾燥 空気、好ましくはアルゴン等の不活性ガスの雰囲気で行 う必要がある。

[0017]

【実施例】以下、実施例にて本発明をより詳細に説明す る。尚、固体電解質のイオン伝導率の測定は次の方法に より行った。固体電解質をフィルム状に成型し、測定用 試料とした。この試料の厚さをマイクロメーターで測定 した後、試料の両面に直径 1 cmの円形プレート状の白金 電極を密若し、この全体を任意の温度に設定できる減圧 容器内に設置し、10-3 Torr以下の高真空まで減圧 して試料の状態が十分に平衡に達した役、LCRメータ ー (横河ヒューレットパッカー F社製4 1 9 2 A) によ れている従来公知の反応手段、例えば、アルカリ等のエ 50 り5 Hz~1 3 MHzの交流電圧を印加し、復素インピーダ

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*4日間真空乾燥したところ。0.3㎜の厚さの透明なフ

ィルムを得た。このフィルムの赤外吸収スペクトルを調

べたところ、水酸基および遊離カルボン酸に由来するビ

ークが認められないこと。エステル生成によるカルボニ ル基の伸縮振動ビークが1740g 「にみられること、

カルボキシレートイオンによる非対称伸縮振動ビークが

1600cm はみられることから、エステル化反応はほぼ完全に行われていることがわかった。フィルムのイオ

ン導電率を測定したところ、25℃で2.0×10°'S

・ローの値が得られた。

化合物(1):

ンス法により伝導率を測定した。

[0018]

【実施例1】固体電解質として本発明によるもの(試料1) および特性比較のため、従来技術によるもの(試料2)を作製した。

[0019]試料1

下記に示される化合物(1)0.384g. 化合物(2)0.450g. 化合物(3)0.167g. 水酸化リチウム12.2mgはよび水0.16gとを撹拌混合し、超音波を照射して十分に溶解させた。この溶液を3cm四 10方のテフロン製の皿に流し込み、ホットプレート上で120℃で2時間加熱した後、真空乾燥器にて140℃で*

化合物(2):

化合物(3):HO(CH,CH,O),,CH, [0020]試料2

試料1で使用した化合物(1)0.278g.化合物(2)0.527g.化合物(3)0.195g および過塩素酸リチウム30.8mqとを撹拌混合し、超音波を照射して十分に溶解せしめ、0.1規定塩酸エタノール溶液3μ1を加えた後、試料1と同様な加熱乾燥処理を行ったところ、0.3mmの厚さの透明なフィルムを得た。このフィルムの赤外吸収スペクトルを調べたところ、水酸基および遊離カルボン酸に由来するビークが認められないこと、エステル生成によるカルボニル基の伸縮振動ピークが1740cm¹にみられることことから、エステル化反応はほぼ完全に行われていることがわかった。イオン導電率を測定したところ、25℃で4.8×10⁻¹S・cm¹の値が得られた。

【0021】直流電圧印加試験

厚さ0.3 mの試料1および試料2の両面に直径1 m. 厚さ0.14mのリチウム笛を密着し、この全体を減圧容器内に設置し、25℃にて10°、Torr以下の高真空まで減圧して試料の状態が十分に平衡に達した後、両リチウム箔に1Vの直流電圧を印加し、流れる電流の経時変化を観察した。試料1では初期の電流5.2μΑが約2日間持続し、このイオン導電性材料が正イオン単独伝導型イオン導電性材料であることが確かめられた。これに対し、試料2では初期の電流79μΑが2日後には50

4、4 u Aまで低下した。次いで試料1または2の固体 電解質を用いてリチウム電池を構成した。正極材料とし て用いたリチウムーマンガン複合酸化物は、飽間らによ ってSanyoTechnical Review,20,114(1989)に述べられた 30 方法により調製した。水酸化リチウム 0.1188と化 学合成二酸化マンガン1gをよく混合した後、375℃ で20時間空気中で加熱し目的物を得た。電極に成型す る際には、このリチウムーマンガン複合酸化物30mgと 導電剤としてのアセチレンブラック20mg、結若剤とし てテフロン5mを加え加圧成型して得た。かかる正極を 用いて作製した電池の断面図を図1に示した。試料1ま たは2の組み込まれたリチウム電池について充放電サイ クルテストを行った。定電流(3.77あるいは37.7 μA/cm¹) で充電時の上限電圧を3.5 V. 放電時の 40 下限電圧を2.5 Vとして行った。図2に代表的なサイ クルにおける常圧-時間曲線を示した。本発明で得られ た試料】を組み込んだ電池(8)では、充放電の繰り返 しにともなう効率の低下はほとんど見られないばかり か、さらに上昇する傾向にある。従来技術で得られた試 料2を組み込んだ電池(b)(比較例 1)では充放電の 繰り返しによる効率の低下がみられた。 図3にはサイク ル毎の放電容量の推移を示した。サイクル毎の放電容量 は、試料1を組み込んだ電池(a)ではあまり変動せず 安定しているのに対し、試料2を組み込んだ電池(h) (比較例1)では安定性に欠け、次第に放電容量も低下

http://www6.ipdl.jpo.go.jp/tjcontenttrns.ipdl?N0000=21&N04.../;%3f%3a%3f%3c9;;%3e///// 6/3/2003

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した。 【0022】

【実施例2】下記に示される化合物(4)108を約500mのトルエン/nーブチルアルコール(1/1)混合溶媒に溶解させ、76、42mの水酸化リチウムを加え、十分に製拌し、化合物(4)中のカルボキシル基を完全にリチウムカルボキシレート化させた。しかる後に溶媒を完全に取り除き、得られた油状物0.772gと下記に示される化合物(5)0.154gおよび化合物(6)0.074gとを観拌混合し、超音波を照射して十分に溶解させた。この混合物に光増感剤として(4ーイソブロビル)フェニル-1-ハイドロキシイソブロビルケトンを約20m加えた後、3cm方のテフロン製の皿*

*に流し込み、垂直上方より高圧水銀ランブからの160 W/cmの紫外線を5 cmの距離で6 秒照射したところ、0.3 mmの厚さの透明なフィルムを得た。さらにこれを70℃で2 日間減圧乾燥させた後、イオン伝導率を測定したところ、25℃で1.6×10-75・cm7の値が得られた。また、実施例1と同様な直流電圧印加試験を行ったところ、4.7 μ Aの電流が約2 日間持続した。さらに実施例1と同様のリチウム電池を作製したところ、初期の放電容量は2.5 m A h / g であり、20サイク10 ル目には2.3 m A h / g となった。

化合物(4):

[(14]

 $CH_2 = CH - CH_2O (CH_2CH_2O) ... CH_3 - CH = CH_3$

化合物 (6):

CH₂=CH-CH₂O (CH₂CH₂O),₂CH₃ [0023]

【実施例3】下記に示される化合物(7)10gを約500mのトルエン/nープチルアルコール(1/1)混合溶媒に溶解させ、86.67mの水酸化リチウムを加え、十分に機拌し、化合物(7)中のカルボキシル基を完全にリチウムカルボキシレート化させた。しかる後に溶媒を完全に取り除き、得られた油状物0.749gと実施例1で使用した化合物(5)0.170gおよび化合物(6)0.081gとを機拌混合し、超音波を照射して十分に溶解させた。この混合物にヒドロシリル化触媒として2重量%塩化白金酸6水和物(H,PtC1,**

※6H₂O) イソプロビルアルコール溶液を2.47μl加えた後、3 cm四方のテフロン製の皿に流し込み、70℃に調整されたオーブン内で2時間加熱したところ。0.3 cmの厚さの透明なフィルムを得た。さらにこれを70℃で2日間減圧乾燥させた後、イオン伝導率を測定したところ、25℃で1.8×10⁻'S・cm゚の値が得られた。また、実施例1と同様な直流電圧印加試験を行ったところ、4.9μAの電流が約2日間持続した。さらに30 実施例1と同様のリチウム電池を作製したところ。初期の放電容量は3.5 mAh/gであり、20サイクル目には2.9 mAh/gとなった。

[0024]

【発明の効果】本発明のリチウム電池は金属リチウム、リチウム合金もしくはリチウム挿入型炭素を活物質とする負価と、正価とが電解質を介して構成されているリチウム電池において、前記電解質が、炭化水素基を介してケイ素原子に結合した。式、一COO¹Li¹で示されるリチウムカルボキシレート基を含有するボリオルガノシロキサン鎖とボリオキンアルキレン鎖とから構成された高分子共食合体架積物からなるので、安定した電圧と電流を与え、高い信頼性を有するという特徴がある。

(CH₂) ₂O (CH₂CH₂O) ₁₂CH₂

【図面の簡単な説明】

化合物(7):

【図1】図1は本発明のリチウム電池の断面図である。 【図2】図2 a は実施例1で得られたリチウム電池の定電流(3.77μA/cd)下での充放電による電圧の時間的変化を示し、図2 b は比較例1の定電流(37.7μA/cd)下での充放電による電圧の時間変化を示したものである。尚、図2 a および図2 b において v は電圧であり、1 は経過時間である。

【図3】図3aは実施例1で得られたリチウム電池の定 50 電流(3.77µA/cd)下での充放電による放電容量

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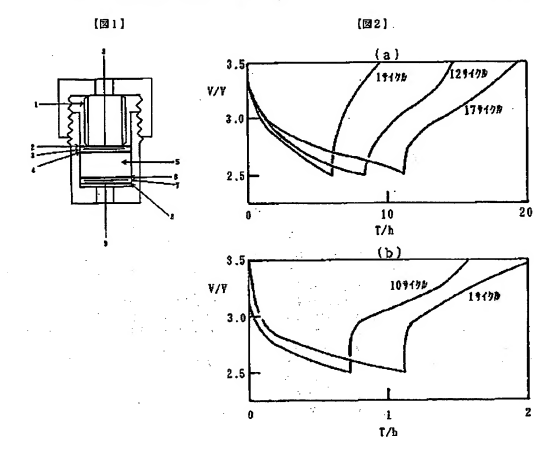
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のサイクル毎の推移を示し、図3 bは比較例1のリチウム電池の定電流 (37.7 μA/cd) 下での充放電による放電容量のサイクル毎の推移を示したものである。 尚、図3 a および図3 bにおいてD.C.は放電容量であり、C.T.はサイクル数である。1はテフロン製スペー*

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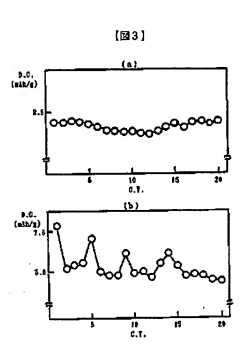
*サー、2はニッケル網,3はニッケル網,4は正額(リチウムーマンガン複合酸化物),5は固体電解質、6は 負額(リチウム箱),7はニッケル網(リチウム箱を圧 着したもの)、8はニッケル網,9はリード線



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CLAIMS

[Claim(s)]

[Claim 1] The formula, -COO-Li+ which the aforementioned electrolyte combined with the silicon atom through the hydrocarbon group in the lithium cell with which the negative electrode which makes an active material a metal lithium, a lithium alloy, or lithium insertion type carbon, and the positive electrode are constituted through the electrolyte Lithium cell characterized by the bird clapper from the macromolecule copolymer bridge formation object which consisted of a polyorganosiloxane chain containing the lithium carboxylate machine shown, and a polyoxyalkylene chain content high molecular compound.

[Claim 2] The lithium cell according to claim 1 manufactured by making the polyoxyalkylene chain content high molecular compound with which a macromolecule copolymer bridge formation object has at least two hydroxyl groups in the polyorganosiloxane which has at least two carboxyl group content hydrocarbon groups in (A)1 molecule, and (B)1 molecule, and the mixture which it becomes from the alkali compound containing (C) lithium metal atom harden by the dehydration condensation reaction.

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DETAILED DESCRIPTION

[Detailed Description of the Invention]

[Industrial Application] this invention relates to a lithium cell, the voltage stabilized by all solid-states in detail, and the lithium cell which gives current.
[0002]

[Description of the Prior Art] In recent years, as for the lithium cell, highly-efficient-izing, a miniaturization, and thin shape-ization are progressing much more with development of microelectronics. Improvement of the electrolyte with which the material of a negative electrode and a positive electrode and improvement of a gestalt are also included in it with last thing in this lithium cell is demanded, and a solid electrolyte is becoming important in the meaning which especially gives a solid state, high flexibility, and high fabricating-operation nature. Furthermore, various advanced properties, such as high ion conductivity, the high lithium ion transference number, high-reliability. and moisture resistance, are demanded. The solid electrolyte material which dissolved lithium salt in the macromolecule resin matrix, and it was made to distribute as such an electrolyte conventionally is known. Composition of the solid electrolyte which combined the polyorganosiloxane chain and the polyethylene-oxide (PEO) chain especially is tried briskly. For example, make a siloxane and PEO construct a bridge over JP,62-209169, A by the hydrosilylation reaction and radiation (electron ray etc.) irradiation by the platinum catalyst, the obtained bridge formation object is made to distribute a lithium ion, and what was made into the solid electrolyte is indicated. Moreover, the lithium cell incorporating these solid electrolytes is indicated by JP,63-170857, A. However, by these methods, in order to dissolve two or more sorts of raw materials, the organic solvent needed to be used. Use of the organic solvent had a possibility of having caused complicated-ization of about [being inapplicable to some products] or a process, and causing aggravation of a work environment, the injury on the charge of an edge strip, remains of the organic solvent to an end product, etc. Moreover, even if it could dissolve the raw material by the organic solvent, the conclusion nature of crosslinking reaction was not completely guaranteed by starting phase separation in process in which the organic solvent evaporates etc., but since it was connected with a lack [deterioration of the quality of a product, and repeatability] etc. after all, there was a problem in utilization. Furthermore, the lithium cell which included the solid electrolyte which made the polystyrene which carried out the graft of the polyethylene-oxide chain distribute lithium salt in JP,2-230667,A is indicated. Although the homogeneity of material improved in this method, there were the following faults. That is, in the solid electrolyte of form which distributes the lithium salt stated by JP,63-170857,A, JP,2-230667,A, etc., it was that to which movement of an anion takes place simultaneously with a lithium ion by electric conduction. In many cases, movement of an anion has many evils and bird clappers. For example, when polarization happens into an electrolyte by movement of an anion at the time of charge and discharge, the structure of an electrode / electrolyte interface will change with time, and the stationary current accompanying movement of a metal ion will not flow. Although an anion cannot but exist for charge compensation with a cation, production of reducing the mobility if possible, i.e., a cation independent conduction-type solid electrolyte, is tried. For example, after making Chemical Society of Japan 1988 spring annual convention lecture number 2XIIC08 carry out the plasma polymerization of the benzenesulfonic-acid methyl ester to octamethylcyclotetrasiloxane. it is made to compound PEO and the lithium ion independent conduction-type solid electrolyte thin http://www4.ipdl.jpo.go.jp/cgi-bin/tran web cgi ejje 6/3/2003

film to which sulfonate ion was fixed is indicated by processing with an iodation lithium further. However, the plasma polymerization had the fault of it having been difficult to acquire the structure of the macromolecule which a reaction means [complicated hatchet], and not becoming a perfect lithium ion independent conduction-type solid electrolyte, in this case. Furthermore, there was a fault that the application range had restrictions, for a plasma polymerization. Thus, even if it was the case where a solid electrolyte was incorporated even if, since the lithium cell proposed till present all had the fault in the property or the manufacture method of the solid electrolyte itself, it was not what the property and manufacturing method of a cell should satisfy.

[0003] When there were no above faults, the material which makes it come to distribute a lithium ion in a specific macromolecule copolymer bridge formation object was excellent in ion conductivity, and may have turned into cation independent conduction-type solid electrolyte material by moreover operating composition, as a result of inquiring wholeheartedly that this trouble should be solved, and it incorporated as an electrolyte of a lithium cell, this invention persons found out that the above faults could be wiped away, and reached this invention. The purpose of this invention is to offer all the solid-state lithium cells that give the stable voltage and current.

[The solution means and operation of a technical problem] This this invention is the formula and - COO-Li+ which the aforementioned electrolyte combined with the silicon atom through the hydrocarbon group in the lithium cell with which the negative electrode which makes an active material a metal lithium, a lithium alloy, or lithium insertion type carbon, and the positive electrode are constituted through the electrolyte. It is related with the lithium cell characterized by the bird clapper from the macromolecule copolymer bridge formation object which consisted of a polyorganosiloxane chain containing the lithium carboxylate machine shown, and a polyoxyalkylene chain content high molecular compound.

[0005] The lithium cell as used in the field of this invention points out both the primary cell which are all solid-states, and the rechargeable battery in which charge and discharge are possible for explaining this. As a factor which determines primary or secondary, it is mainly the kind of positive active material. In this invention, since it aims at the lithium ion independent conductivity previously described as one of the highly-efficient-izing of a lithium cell, it is necessary to make second order so-called rocking-chair type composition also in which the first order as a form of a lithium cell. That is, managing the cell reaction in positive/negative two poles is taken only as a lithium ion. Therefore, it is necessary to be the form from which it sets to a positive electrode, and a lithium ion is inserted and (at the time of electric discharge) desorbed (at the time of the charge in a rechargeable battery). Although it does not have the sandwich structure of a negative-electrode-solid electrolyte-positive electrode as structure of a cell and a size, thickness, and a gestalt are not limited, Oshi's thing is typically possible even for a coin type and the cylinder type of a spiral method from area 2 and the paper type with a thickness of about about 1mm of several cm. Although a negative electrode uses a lithium metallic foil typically, especially if the lithium metal as a negative-electrode active material is the conductive sheet which exists with the gestalt effective in electrode reaction, it will not be limited. For example, the thing by which the lithium etc. was held, or a lithium-aluminium alloy is mentioned to a carbon sheet. Although it will not be limited as an active material of a positive electrode especially if the above-mentioned cell reaction can be started, an inorganic stratified compound is used typically. For example, manganese dioxide, oxidization vanadium, 2 titanium sulfides, cobalt oxide, nickel oxide, molybdenum sulfides, these complex, etc. are used. It is usually weak, and the work to which conductivity uses an organic resin for a low sake as caking additive, and it is made a tabular is carried out, or, as for these compounds, a carbon particle etc. is added as a conductive grant agent. Moreover, it is suitable if the stratified compound which made structure incorporate a lithium atom in part from the permanence of the layer structure by the cycle of charge and discharge or the absorptivity and desorptivity of a lithium ion is used when building especially a rechargeable battery.

[0006] subsequently, explaining the above-mentioned macromolecule copolymer bridge formation object this bridge formation object -- the alkylene machine of the carbon numbers 1-8, such as a methylene group, an ethylene, a propylene machine, a butylene machine, a pentylene machine, a

hexelene machine, a heptylene machine, and an octylene machine, -- a hydrocarbon group which is illustrated with the arylene machine of the carbon numbers 8-20, such as; phenylene group and a naphthylene machine That what is necessary is just to consist of the lithium carboxylate machine content polyorganosiloxane chains and polyoxyalkylene chain content high molecular compounds which were minded and were combined with the silicon atom and which have a formula and the chemical structure shown by -COO-Li+ It is not limited about the kind of the high molecular compound, a gestalt, and especially crosslinking density.

[0007] Since the basis (-COO-) in which the solid electrolyte of this invention contains the above anions is combined with the silicon atom through the hydrocarbon group, the mobility of an anion is extremely low. Therefore, in this macromolecule copolymer bridge formation object, a lithium ion dissociates from an anion, and when a lithium ion moves, ion conductivity is discovered. If the kind of anion which exists in a macromolecule copolymer bridge formation object is only -COO-, this solid electrolyte material will work as a lithium ion independent conductor.

[0008] The method of the following [method / desirable about the manufacture method of this solid electrolyte] is recommended. "How to manufacture by making the polyoxyalkylene chain content high molecular compound which has at least two hydroxyl groups in the polyorganosiloxane which has at least two carboxyl group content hydrocarbon groups in (A)1 molecule, and (B)1 molecule, and the mixture which consists of an alkali compound containing (C) lithium metal atom harden by the dehydration condensation reaction."

[0009] The ratio of the siloxane unit which it is required to have two or more carboxyl group content hydrocarbon groups, and this carboxyl group content hydrocarbon group combined in 1 molecule in order for the polyorganosiloxane of the (A) component to form a macromolecule copolymer bridge formation object for explaining this, and the other siloxane unit What is within the limits of 0.01-100 is desirable. Although any of the shape of a straight chain, the letter of branching, annular, reticulated, and the three-dimensional structure are sufficient as the molecular structure of this component, as for the ease of formation of a macromolecule copolymer bridge formation object to more than the half, it is desirable that they are the shape of a straight chain and a letter of branching. Moreover, although especially the molecular weight is not limited, in order to obtain the moderate hardness as the ease of manufacture, and a macromolecule copolymer bridge formation object, it is desirable that it is within the limits of 1 million-1 million. (A) As a carboxyl group content hydrocarbon group of a component, the basis shown by general formula HOOC-R1- (R1 is the arylene machine of the carbon numbers 6-20, such as an alkylene machine of the carbon numbers 1-8, such as a methylene group, an ethylene, a propylene machine, a butylene machine, a pentylene machine, a hexylene machine, a heptylene machine, and an octylene machine, or a phenylene group, and a naphthylene machine, among a formula.) is mentioned, for example, also in these, a carboxy (A) As organic machines other than the carboxyl group content hydrocarbon group in a component, aralkyl machines, such as aryl group; benzyls, such as alkyl group; phenyl groups, such as a methyl group, an ethyl group, and a propyl group, a tolyl group, and a xylyl group, and a phenethyl machine, are illustrated, for example. Moreover, as a basis combined with the silicon atom, a small amount of hydrogen atom and an alkoxy group may be contained. As for more than the half of the ORGANO machine combined with the silicon atom from a viewpoint of economical efficiency and the formation nature of a good macromolecule copolymer bridge formation object, it is desirable that it is a methyl group. As this polyorganosiloxane, the methyl carboxy propyl siloxane dimethylsiloxane copolymer by which chain both ends were blocked with the trimethylsiloxy machine, and the methyl carboxy propyl siloxane methylphenyl siloxane copolymer by which chain both ends were blocked with the trimethylsiloxy machine are mentioned, for example. Although various methods are learned as the synthetic method of this polyorganosiloxane, the annular object and end halt agent which are obtained by carrying out the cohydrolysis of the ORGANO dichlorosilane which has a cyano group as the one method, and the ORGANO dichlorosilane which does not have a cyano group are stirred in sulfuric-acid solution, and the reaction which a cyano group converts into a carboxyl group, and the method of making ring opening polymerization start are mentioned.

[0010] (B) The high molecular compound of a component is the cross linking agent of the above-mentioned (A) component, and in order to carry out the work as a cross linking agent, it needs to

contain at least two hydroxyl groups in 1 molecule. Moreover, although it is desirable to have polyoxyalkylene chain in a macromolecule copolymer bridge formation object for a high ion conductivity manifestation, for that, it is required in the high-molecular-compound structure of the (B) component to contain polyoxyalkylene chain.

[0011] By the way, it is very important that the (A) component and the (B) component dissolve mutually in order that using a solvent in process may bring about the conclusion of about [avoid] and crosslinking reaction, the structure of a bridge formation product, and the good repeatability of physical properties. (A) It is desirable to have the siloxane unit about the molecular structure and the chemical structure of the (B) component for dissolving with a component, and it is a general formula [-izing 1] from the ease of formation of a macromolecule copolymer bridge formation object etc.

(-- an organic machine univalent in R2, an organic machine divalent in R3, an organic machine univalent [4/R] in an alkylene machine and R5, and a and c are / integer / of 2-1000 / and p of integer / of 0-1000 / and b / the integers of 2-100 among a formula The polyorganosiloxane which has polyoxyalkylene chain which has a hydroxyl group at the end shown by) as an at least 2 graft chain is desirable. As for this organopolysiloxane, aralkyl machines, such as aryl group; benzyls, such as alkyl group; phenyl groups, such as a methyl group, an ethyl group, and a propyl group, a tolyl group, and a xylyl group, and a phenethyl machine, are illustrated among an upper formula, as for R2. Moreover, a small amount of hydrogen atom and an alkoxy group may be contained in part. As for more than the half of R2, from a viewpoint of economical efficiency and the formation nature of a good macromolecule copolymer bridge formation object, it is desirable that it is a methyl group. R3 is divalent organic machines, such as arylene machines, such as an alkylene machine of the carbon numbers 1-8, such as a methylene group, an ethylene, a propylene machine, a butylene machine, a pentylene machine, a hexylene machine, a heptylene machine, and an octylene machine, or a phenylene group, and a naphthylene machine. R4 is alkylene machines, such as a methylene group, an ethylene, a propylene machine, a butylene machine, a pentylene machine, a hexylene machine, and a heptylene machine. R5 is acyl groups, such as an alkyl group; acetyl group or propionyl machines, such as a methyl group, an ethyl group, and a propyl group. Although a and c are within the limits of 0-1000, b is within the limits of 2-1000 and it is not limited especially, respectively, in order for the (A) component and the (B) component to dissolve, a and the ratio of (b+c) have desirable within the limits of -(1:5)(5:1).

[0012] Although various methods are learned as the synthetic method of this graft copolymer The polyoxyalkylene to which a part of side chain has an unsaturation hydrocarbon group at the piece end, and it has an acyloxy machine at the other ends as the one method at the polyorganosiloxane which replaced by the hydrogen atom, The graft of the polyoxyalkylene which has an unsaturation hydrocarbon group at the piece end, and has a trimethylsilyl machine at the other ends is carried out by the hydrosilylation reaction with a predetermined ratio. The method of converting only the trimethylsilyl machine of a graft end into a hydroxyl group by the alcohol of an excessive amount is mentioned to after an appropriate time.

[0013] (B) Although a component is a high molecular compound which has at least two hydroxyl groups in the above 1 molecules, and contains polyoxyalkylene chain in structure When higher ion conductivity is required, this high molecular compound is general formula HO-(R6O) q-H (an alkylene machine and q of R6 are the integers of 1-100 among a formula.). Polyoxyalkylene or HO-(R7O) r-R8 (an organic machine univalent [7/R] in an alkylene machine and R8 and r are the integers of 1-100 among a formula.) which has a hydroxyl group in the chain both ends shown It is desirable that it is polyoxyalkylene which has a hydroxyl group at the piece end of a chain shown. R6 and R7 of the upper formula of this polyoxyalkylene are alkylene machines, such as a methylene group, an ethylene, a propylene machine, a butylene machine, a pentylene machine, a hexylene

machine, and a heptylene machine. R7 is acyl groups, such as an alkyl group; acetyl group or propionyl machines, such as a methyl group, an ethyl group, and a propyl group. q and r are within the limits of 1-100, and its within the limits of 5-20 is desirable. The condensation reaction of the polyoxyalkylene which has a hydroxyl group at the polyoxyalkylene or the piece end of a chain which has a hydroxyl group in these chain both ends is carried out to the above-mentioned (A) component, and it carries out the work which increases the content of the bridge formation chain of polyoxyalkylene, or a graft chain into a macromolecule copolymer bridge formation object. Even if this bridge formation chain or a graft chain does not exist, the bridge formation chain or graft chain of polyoxyalkylene can be made to form by the polyoxyalkylene chain in a high molecular compound in this invention. Since there is a limitation in the amount of introduction of the polyoxyalkylene chain in a high molecular compound in the conditions which the (A) component and the (B) component dissolve, make it however, more desirable to contain the polyoxyalkylene which has a hydroxyl group at the polyoxyalkylene or the piece end of a chain which has a hydroxyl group in chain both ends in the (B) component. Moreover, in this invention, the direction where the graft chain of polyoxyalkylene existed a little in the macromolecule copolymer bridge formation object is in the inclination whose ion conductivity improves.

[0014] (C) Although a component is an alkali compound containing a lithium atom and especially the kind is not limited, a hydroxide, an alcoholate, a hydride, etc. are desirable and it is desirable to be used, especially the gestalt of a hydroxide, i.e., a lithium hydroxide, (LiOH). This (C) component carries out the role which finally forms the carboxyl group in the (A) component into lithium carboxylate by dehydration at the same time it carries out the catalysis of the esterification reaction of the (A) component and the (B) component. Consequently, as a macromolecule copolymer bridge formation object, it is fixed to a siloxane polymer chain and an anion (carboxylate ion) becomes the gestalt by which the cation (lithium ion) which is a counter ion was distributed. Moreover, as for the variance, it is desirable that ratio ratio [of the number of mols [Li+] of a lithium ion to the number of mols of the oxy-alkylene machine in a macromolecule copolymer bridge formation object [RO] [Li+]/[RO] sets to 0.005-0.25, and it is 0.02-0.1 more preferably. this -- this ratio if 0.25 is exceeded -- the polarity of a macromolecule copolymer bridge formation object -- going up -- the maneuverability of a segment -- bad -- becoming -- moreover -- It is because ion conductivity high from the fall of the number of carriers will become difficult to get if it becomes less than 0.005. Moreover, the number of mols of the carboxyl group in the (A) component, the number of mols of the hydroxyl group in the (B) component, and the ratio of the sum total of the alkali equivalent in the (C) component are within the limits of - (1:10) (10:1), and its within the limits of - (1.0:1.2) (1.2:1.0) is desirable. Although the macromolecule copolymer bridge-formation object which the esterification reaction occurred and solidified by any ratio is obtained, this has (A) - (C) component in the inclination which will be in the state where of an unreacted carboxyl group or an unreacted hydroxyl group remained in solid electrolyte material, when the ratios of the number of mols of the number of mols of the hydroxyl group in the (B) component, the sum total of the alkali equivalent in the (C) component, and the carboxyl group in the (A) component differ sharply. Since it is expected that evils, such as a reaction with electrode material, happen when the solid electrolyte in this state is included in a cell, it is not desirable. Moreover, if the above-mentioned mole ratio is set to 1, an anion kind will serve as only carboxylate ion to which only the lithium ion was fixed, and a cation kind will serve as a perfect lithium ion independent conduction-type solid electrolyte.

[0015] Although the mixture which consists of a (A) component - (C) component in this method is stiffened by the dehydration condensation reaction and a macromolecule copolymer bridge formation object is made to form, this bridge formation object is mainly formed of the esterification reaction of the carboxyl group in the (A) component, and the hydroxyl group of the (B) component. Esterification reaction catalysts, such as a conventionally well-known reaction means, for example, alkali etc., by which it is used for the reaction of a carboxyl group and a hydroxyl group as the reaction means, are used, and the method of making it react to the bottom of ordinary temperature or heating can be used. Moreover, in order that heating may remove the water which is the by-product generated by the esterification reaction at the same time it promotes this esterification reaction, it is an effective technical means, and the temperature is usually 150 degrees C or less. Moreover, this

crosslinking reaction can be performed on condition that a non-solvent. (A) A component and the (B) component are uniformly mixable easily by operation of stirring etc. (C) You may add, after making it dissolve in the (B) component beforehand or mixing the (A) component and the (B) component, since a component has solubility to an oxy-alkylene chain. (C) Although operation of stirring etc. is also possible for the dissolution to the oxy-alkylene chain of a component, for dissolution time shortening, operation of heating or ultrasonic irradiation or addition of very little water is effective. Moreover, when use of the organic solvent is permissible in a dissolution process, a (A) component -(C) component may be mixed and dissolved in the organic solvent, and a solvent may be evaporated after an appropriate time. Although especially this organic solvent is not limited, a tetrahydrofuran, a dioxane, an acetonitrile, a JIMECHI formamide, and dimethyl sulfoxide are mentioned, for example. Moreover, since this esterification reaction generates water as a by-product, it is desirable to put the mixture of a (A) component - (C) component under reduced pressure finally, for example, the method of heating under reduced pressure, after advancing an esterification reaction to some extent under ordinary-pressure heating, and performing simultaneously the removal of water and the conclusion of an esterification reaction which were generated is recommended. However, when a solvent is used for a dissolution process, after advancing an esterification reaction to some extent below with the boiling point of the organic solvent under an ordinary pressure, the organic solvent is evaporated and it is necessary to heat under reduced pressure after an appropriate time.

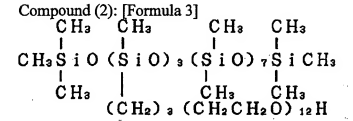
[0016] Although the lithium cell of this invention comes to combine the above-mentioned positive electrode, a negative electrode, and a solid electrolyte, it is not limited especially about the building method. You may combine three persons, after producing each separately. However, since the efficiency as a cell increases so that the touch area in an interface is generally large, after carrying out the cast of the mixture of a solid electrolyte raw material on a positive electrode, making crosslinking reaction perform and film-izing, the method of making a negative electrode rival is recommended. in order that [moreover,] a negative electrode and a solid electrolyte may receive transformation with moisture -- production of a cell -- dry air -- it is necessary to carry out in the atmosphere of inert gas, such as an argon, preferably

[0017].

[Example] Hereafter, an example explains this invention to a detail more. In addition, measurement of the ionic conductivity of a solid electrolyte was performed by the following method. The solid electrolyte was cast in the shape of a film, and it considered as test sample. After measuring the thickness of this sample with a micrometer, the platinum electrode of the shape of a circular plate with a diameter of 1cm is stuck to both sides of a sample. After installing in the reduced pressure container which can set this whole as arbitrary temperature, decompressing to the high vacuum of 10 to 5 or less Torrs and the state of a sample fully reaching a balance, 5Hz - 13MHz alternating voltage was impressed by the LCR meter (4192made from YOKOGAWA ELECTRIC Hewlett Packard A), and conductivity was measured by the complex impedance method.

[Example 1] What is depended on the conventional technology (sample 2) was produced for the thing (sample 1) depended on this invention as a solid electrolyte, and property comparison.

[0019] Stirring mixture of compound (1)0.384g, compound (2)0.450g, compound (3)0.167g, 12.2mg of lithium hydroxides, and 0.16g of water shown in the sample 1 following was carried out, the ultrasonic wave was irradiated, and it was made to fully dissolve. After slushing this solution into the pan made from the Teflon of 3cm around and heating it at 120 degrees C on a hot plate for 2 hours, when the vacuum drying was carried out for four days at 140 degrees C by the vacuum dryer, the film with the transparent thickness of 0.3mm was obtained. Since that the peak originating in a hydroxyl group and an isolation carboxylic acid is not accepted, the stretching-vibration peak of the carbonyl group by esterification being seen by 1740cm-1, and the unsymmetrical stretching-vibration peak by carboxylate ion were seen by 1600cm-1 when the infrared absorption spectrum of this film was investigated, it turns out that the esterification reaction is performed nearly completely. When the ion conductivity of a film was measured, the value of 2.0x10-7 S-cm -1 was acquired at 25 degrees C. Compound (1): [Formula 2]



Compound (3): HO(CH2CH2O)12CH3 [0020] After having carried out stirring mixture of compound (1)0.278g, compound (2)0.527g, compound (3)0.195g, and 30.8mg of lithium perchlorates used by sample 2 sample 1, having irradiated the ultrasonic wave, making it fully dissolve and adding 3micro of decinormal hydrochloric-acid ethanol solutions l, when the same stoving processing as a sample 1 was performed, the film with the transparent thickness of 0.3mm was obtained. When the infrared absorption spectrum of this film was investigated, it turns out [that the peak originating in a hydroxyl group and an isolation carboxylic acid is not accepted, and / as which the stretching-vibration peak of the carbonyl group by esterification is regarded by 1740cm-1] that the shell and the esterification reaction are rattlingly performed nearly completely. When ion conductivity was measured, the value of 4.8x10-5 S-cm -1 was acquired at 25 degrees C.

[0021] The lithium foil with a diameter [of 1cm] and a thickness of 0.14mm was stuck to both sides of the sample 1 with a direct-current-voltage impression examination thickness of 0.3mm and a sample 2, this whole was installed in the reduced pressure container, after it decompressed to the high vacuum of 10 to 5 or less Torrs at 25 degrees C and the state of a sample fully reached the balance, the direct current voltage of 1V was impressed to both the lithiums foil, and flowing aging of current was observed. By the sample 1, 5.2micro of early current A continued for about two days, and it was confirmed that this ion conductivity material is cation independent conduction-type ion conductivity material. On the other hand, two days after by the sample 2, 79micro of early current A fell to 4.4microA. Subsequently, the lithium cell was constituted using the solid electrolyte of samples 1 or 2. The lithium-manganese multiple oxide used as a positive-electrode material was prepared by the method stated to Sanyo Technical Review and 20,114 (1989) by Noma and others. Lithium hydroxide After often mixing 0.118g and 1g of chemosynthesis manganese dioxide, it heated in 20-hour air at 375 degrees C, and the specified substance was obtained. When casting to an electrode, Teflon 5mg was added as acetylene black 20mg as 30mg of this lithium-manganese multiple oxide, and an electric conduction agent, and a binder, pressurization molding was carried out, and it obtained. The cross section of the cell produced using this positive electrode was shown in drawing 1. The chargeand-discharge cycle test was performed about the lithium cell with which samples 1 or 2 were incorporated. 3.5V and minimum voltage at the time of electric discharge were performed for the upper limit voltage at the time of charge as 2.5V by the constant current (3.77 or 37.7microA/cm -2). The voltage-time curve in a cycle typical to drawing 2 was shown. By the cell (a) incorporating the sample 1 obtained by this invention, not seeing and the inclination to go up further has most decline in the efficiency accompanying the repeat of charge and discharge. In the cell (b) incorporating the sample 2 obtained with the conventional technology, and the (example 1 of comparison), decline in the efficiency by the repeat of charge and discharge was seen. Transition of the service capacity for every cycle was shown in drawing 3. To having seldom changed the service capacity for every cycle by the cell (a) incorporating the sample 1, but being stable, in the cell (b) incorporating the sample 2, and the (example 1 of comparison), stability was missing and service capacity also fell gradually. **[0022]**

[Example 2] Compound (4)10g shown below was dissolved in about 500ml toluene / n-butyl alcohol (1/1) mixed solvent, the 76.42mg lithium hydroxide was added, it fully stirred, and the carboxyl http://www4.ipdl.jpo.go.jp/cgi-bin/tran web cgi ejje 6/3/2003

group in a compound (4) was made to form into lithium carboxylate completely. Stirring mixture of compound (5)0.154g which removes a solvent completely after an appropriate time, and is shown in 0.772g of obtained oily matter and the following, and the compound (6)0.074g was carried out, the ultrasonic wave was irradiated, and it was made to fully dissolve. After adding about 20mg of phenyl-1-hide ROKISHI isopropyl ketones to this mixture as a photosensitizer (4-isopropyl), when it slushed into the pan made from the Teflon of 3cm around and the ultraviolet rays of 160 W/cm from a high-pressure mercury lamp were irradiated for 6 seconds in 5cm distance from the perpendicular upper part, the film with the transparent thickness of 0.3mm was obtained. After carrying out reduced pressure drying of this for two days at 70 degrees C furthermore, when ionic conductivity was measured, the value of 1.6x10-7 S-cm -1 was acquired at 25 degrees C. Moreover, when the same direct-current-voltage impression examination as an example 1 was performed, the current of 4.7microA continued for about two days. When the still more nearly same lithium cell as an example 1 was produced, early service capacity is 2.5 mAh/g and became 20 cycle eye with 2.3 mAh/g.

Compound (5): CH2=CH-CH2O(CH2CH2O)12CH2-CH=CH2 compound (6):CH2=CH-CH2O (CH2CH2O)12CH3 [0023]

[Example 3] Compound (7)10g shown below was dissolved in about 500ml toluene / n-butyl alcohol (1/1) mixed solvent, the 86.67mg lithium hydroxide was added, it fully stirred, and the carboxyl group in a compound (7) was made to form into lithium carboxylate completely. Stirring mixture of compound (5)0.170g which removed the solvent after an appropriate time completely, and used it for it in 0.749g of obtained oily matter and the example 1, and the compound (6)0.081g was carried out, the ultrasonic wave was irradiated, and it was made to fully dissolve. After 2.47microl Adding a chloroplatinic-acid 6 hydrate (H2PtCl6.6H2O) isopropyl alcohol solution to this mixture 2% of the weight as a hydrosilylation catalyst, it slushed into the pan made from the Teflon of 3cm around, and when heated within the oven adjusted to 70 degrees C for 2 hours, the film with the transparent thickness of 0.3mm was obtained. After carrying out reduced pressure drying of this for two days at 70 degrees C furthermore, when ionic conductivity was measured, the value of 1.8x10-7 S-cm -1 was acquired at 25 degrees C. Moreover, when the same direct-current-voltage impression examination as an example 1 was performed, the current of 4.9microA continued for about two days. When the still more nearly same lithium cell as an example 1 was produced, early service capacity is 3.5 mAh/g and became 20 cycle eye with 2.9 mAh/g.

[0024]

[Effect of the Invention] In the lithium cell with which the negative electrode to which the lithium cell of this invention makes an active material a metal lithium, a lithium alloy, or lithium insertion type carbon, and the positive electrode are constituted through the electrolyte The aforementioned electrolyte combined with the silicon atom through the hydrocarbon group. A formula, -COO-Li+Since it consists of a macromolecule copolymer bridge formation object which consisted of a polyorganosiloxane chain containing the lithium carboxylate machine shown, and polyoxyalkylene http://www4.ipdl.jpo.go.jp/cgi-bin/tran web cgi ejje

| chain The stable voltage and stable current are given and there is the feature of having high rel | iability. |
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TECHNICAL FIELD

[Industrial Application] this invention relates to a lithium cell, the voltage stabilized by all solid-states in detail, and the lithium cell which gives current.

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EFFECT OF THE INVENTION

[Effect of the Invention] In the lithium cell with which the negative electrode to which the lithium cell of this invention makes an active material a metal lithium, a lithium alloy, or lithium insertion type carbon, and the positive electrode are constituted through the electrolyte The aforementioned electrolyte combined with the silicon atom through the hydrocarbon group. A formula, -COO-Li+Since it consists of a macromolecule copolymer bridge formation object which consisted of a polyorganosiloxane chain containing the lithium carboxylate machine shown, and polyoxyalkylene chain, the stable voltage and stable current are given and there is the feature of having high reliability.

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TECHNICAL PROBLEM

[Description of the Prior Art] In recent years, as for the lithium cell, highly-efficient-izing, a miniaturization, and thin shape-ization are progressing much more with development of microelectronics. Improvement of the electrolyte with which the material of a negative electrode and a positive electrode and improvement of a form are also included in it with last thing in this lithium cell is demanded, and a solid electrolyte is becoming important in the meaning which especially gives a solid state, high flexibility, and high fabricating-operation nature. Furthermore, various advanced properties, such as high ion conductivity, the high lithium ion transference number, high-reliability, and moisture resistance, are demanded. The solid electrolyte material which dissolved lithium salt in the macromolecule resin matrix, and it was made to distribute as such an electrolyte conventionally is known. Composition of the solid electrolyte which combined the polyorganosiloxane chain and the polyethylene-oxide (PEO) chain especially is tried briskly. For example, make a siloxane and PEO construct a bridge over JP,62-209169,A by the hydrosilylation reaction and radiation (electron ray etc.) irradiation by the platinum catalyst, the obtained bridge formation object is made to distribute a lithium ion, and what was made into the solid electrolyte is indicated. Moreover, the lithium cell incorporating these solid electrolytes is indicated by JP,63-170857, A. However, by these methods, in order to dissolve two or more sorts of raw materials, the organic solvent needed to be used. Use of the organic solvent had a possibility of having caused complicated-ization of about [being inapplicable to some products] or a process, and causing aggravation of a work environment, the injury on the charge of an edge strip, remains of the organic solvent to an end product, etc. Moreover, even if it could dissolve the raw material by the organic solvent, the conclusion nature of crosslinking reaction was not completely guaranteed by starting phase separation in process in which the organic solvent evaporates etc., but since it was connected with a lack [deterioration of the quality of a product, and repeatability] etc. after all, there was a problem in utilization. Furthermore, the lithium cell which included the solid electrolyte which made the polystyrene which carried out the graft of the polyethylene-oxide chain distribute lithium salt in JP,2-230667,A is indicated. Although the homogeneity of material improved in this method, there were the following faults. That is, in the solid electrolyte of form which distributes the lithium salt stated by JP,63-170857,A, JP,2-230667,A, etc., it was that to which movement of an anion takes place simultaneously with a lithium ion by electric conduction. In many cases, movement of an anion has many evils and bird clappers. For example, when polarization happens into an electrolyte by movement of an anion at the time of charge and discharge, the structure of an electrode / electrolyte interface will change with time, and the stationary current accompanying movement of a metal ion will not flow. Although an anion cannot but exist for charge compensation with a cation, production of reducing the mobility if possible, i.e., a cation independent conduction-type solid electrolyte, is tried. For example, after making Chemical Society of Japan 1988 spring annual convention lecture number 2XIIC08 carry out the plasma polymerization of the benzenesulfonic-acid methyl ester to octamethylcyclotetrasiloxane, it is made to compound PEO and the lithium ion independent conduction-type solid electrolyte thin film to which sulfonate ion was fixed is indicated by processing with an iodation lithium further. However, the plasma polymerization had the fault of it having been difficult to acquire the structure of the macromolecule which a reaction means [complicated hatchet], and not becoming a perfect lithium ion independent conduction-type solid electrolyte, in this case. Furthermore, there was a fault that the application range had restrictions, for a plasma polymerization. Thus, even if it was the case http://www4.ipdl.jpo.go.jp/cgi-bin/tran web_cgi ejje 6/3/2003

where a solid electrolyte was incorporated even if, since the lithium cell proposed till present all had the fault in the property or the manufacture method of the solid electrolyte itself, it was not what the property and manufacturing method of a cell should satisfy.

[0003] When there were no above faults, the material which makes it come to distribute a lithium ion in a specific macromolecule copolymer bridge formation object was excellent in ion conductivity, and may have turned into cation independent conduction-type solid electrolyte material by moreover operating composition, as a result of inquiring wholeheartedly that this trouble should be solved, and it incorporated as an electrolyte of a lithium cell, this invention persons found out that the above faults could be wiped away, and reached this invention. The purpose of this invention is to offer all the solid-state lithium cells that give the stable voltage and current.

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OPERATION

[The solution means and operation of a technical problem] In the lithium cell with which the negative electrode to which this this invention makes an active material a metal lithium, a lithium alloy, or lithium insertion type carbon, and the positive electrode are constituted through the electrolyte The formula, -COO-Li+ which the aforementioned electrolyte combined with the silicon atom through the hydrocarbon group It is related with the lithium cell characterized by the bird clapper from the macromolecule copolymer bridge formation object which consisted of a polyorganosiloxane chain containing the lithium carboxylate machine shown, and a polyoxyalkylene chain content high molecular compound.

[0005] The lithium cell as used in the field of this invention points out both the primary cell which are all solid-states, and the rechargeable battery in which charge and discharge are possible for explaining this. As a factor which determines primary or secondary, it is mainly the kind of positive active material. In this invention, since it aims at the lithium ion independent conductivity previously described as one of the highly-efficient-izing of a lithium cell, it is necessary to make second order so-called rocking-chair type composition also in which the first order as a form of a lithium cell. That is, managing the cell reaction in positive/negative two poles is taken only as a lithium ion. Therefore, it is necessary to be the form from which it sets to a positive electrode, and a lithium ion is inserted and (at the time of electric discharge) desorbed (at the time of the charge in a rechargeable battery). Although it does not have the sandwich structure of a negative-electrode-solid electrolyte-positive electrode as structure of a cell and a size, thickness, and a gestalt are not limited, Oshi's thing is typically possible even for a coin type and the cylinder type of a spiral method from area 2 and the paper type with a thickness of about about 1mm of several cm. Although a negative electrode uses a lithium metallic foil typically, especially if the lithium metal as a negative-electrode active material is the conductive sheet which exists with the gestalt effective in electrode reaction, it will not be limited. For example, the thing by which the lithium etc. was held, or a lithium-aluminium alloy is mentioned to a carbon sheet. Although it will not be limited as an active material of a positive electrode especially if the above-mentioned cell reaction can be started, an inorganic stratified compound is used typically. For example, manganese dioxide, oxidization vanadium, 2 titanium sulfides, cobalt oxide, nickel oxide, molybdenum sulfides, these complex, etc. are used. It is usually weak, and the work to which conductivity uses an organic resin for a low sake as caking additive, and it is made a tabular is carried out, or, as for these compounds, a carbon particle etc. is added as a conductive grant agent. Moreover, it is suitable if the stratified compound which made structure incorporate a lithium atom in part from the permanence of the layer structure by the cycle of charge and discharge or the absorptivity and desorptivity of a lithium ion is used when building especially a rechargeable battery.

[0006] subsequently, explaining the above-mentioned macromolecule copolymer bridge formation object this bridge formation object -- the alkylene machine of the carbon numbers 1-8, such as a methylene group, an ethylene, a propylene machine, a butylene machine, a pentylene machine, a hexelene machine, a heptylene machine, and an octylene machine, -- a hydrocarbon group which is illustrated with the arylene machine of the carbon numbers 8-20, such as; phenylene group and a naphthylene machine It consisted of the lithium carboxylate machine content polyorganosiloxane chains and polyoxyalkylene chain content high molecular compounds which were minded and were combined with the silicon atom and which have a formula and the chemical structure shown by -

COO-Li+. About the kind of the high molecular compound, a gestalt, and especially crosslinking density, it is not limited that what is necessary is just to be.

[0007] Since the basis (-COO-) in which the solid electrolyte of this invention contains the above anions is combined with the silicon atom through the hydrocarbon group, the mobility of an anion is a low extremely. Therefore, in this macromolecule copolymer bridge formation object, a lithium ion dissociates from an anion, and when a lithium ion moves, ion conductivity is discovered. If the kind of anion which exists in a macromolecule copolymer bridge formation object is only -COO-, this solid electrolyte material will work as a lithium ion independent conductor.

[0008] The method of the following [method / desirable about the manufacture method of this solid electrolyte] is recommended. "How to manufacture by making the polyoxyalkylene chain content high molecular compound which has at least two hydroxyl groups in the polyorganosiloxane which has at least two carboxyl group content hydrocarbon groups in (A)1 molecule, and (B)1 molecule, and the mixture which consists of an alkali compound containing (C) lithium metal atom harden by the dehydration condensation reaction."

10009] The ratio of the siloxane unit which it is required to have two or more carboxyl group content hydrocarbon groups, and this carboxyl group content hydrocarbon group combined in 1 molecule in order for the polyorganosiloxane of the (A) component to form a macromolecule copolymer bridge formation object for explaining this, and the other siloxane unit What is within the limits of 0.01-100 is desirable. Although any of the shape of a straight chain, the letter of branching, annular, reticulated, and the three-dimensional structure are sufficient as the molecular structure of this component, as for the ease of formation of a macromolecule copolymer bridge formation object to more than the half, it is desirable that they are the shape of a straight chain and a letter of branching. Moreover, although especially the molecular weight is not limited, in order to obtain the moderate hardness as the ease of manufacture, and a macromolecule copolymer bridge formation object, it is desirable that it is within the limits of 1 million-1 million. (A) As the carboxyl group content hydrocarbon group of a component For example, the basis shown by general formula HOOC-R1- (R1 is the arylene machine of the carbon numbers 6-20, such as an alkylene machine of the carbon numbers 1-8, such as a methylene group, an ethylene, a propylene machine, a butylene machine, a pentylene machine, a hexylene machine, a heptylene machine, and an octylene machine, or a phenylene group, and a naphthylene machine, among a formula.) is mentioned, also in these, a carboxy alkyl group is desirable and especially a carboxy propyl group is desirable (A) As organic machines other than the carboxyl group content hydrocarbon group in a component, aralkyl machines, such as aryl group; benzyls, such as alkyl group; phenyl groups, such as a methyl group, an ethyl group, and a propyl group, a tolyl group, and a xylyl group, and a phenethyl machine, are illustrated, for example. Moreover, as a basis combined with the silicon atom, a small amount of hydrogen atom and an alkoxy group may be contained. As for more than the half of the ORGANO machine combined with the silicon atom from a viewpoint of economical efficiency and the formation nature of a good macromolecule copolymer bridge formation object, it is desirable that it is a methyl group. As this polyorganosiloxane, the methyl carboxy propyl siloxane dimethylsiloxane copolymer by which chain both ends were blocked with the trimethylsiloxy machine, and the methyl carboxy propyl siloxane methylphenyl siloxane copolymer by which chain both ends were blocked with the trimethylsiloxy. machine are mentioned, for example. Although various methods are learned as the synthetic method of this polyorganosiloxane, the annular object and end halt agent which are obtained by carrying out the cohydrolysis of the ORGANO dichlorosilane which has a cyano group as the one method, and the ORGANO dichlorosilane which does not have a cyano group are stirred in sulfuric-acid solution, and the reaction which a cyano group converts into a carboxyl group, and the method of making ring opening polymerization start are mentioned.

[0010] (B) The high molecular compound of a component is the cross linking agent of the above-mentioned (A) component, and in order to carry out the work as a cross linking agent, it needs to contain at least two hydroxyl groups in 1 molecule. Moreover, although it is desirable to have polyoxyalkylene chain in a macromolecule copolymer bridge formation object for a high ion conductivity manifestation, for that, it is required in the high-molecular-compound structure of the (B) component to contain polyoxyalkylene chain.

[0011] By the way, it is very important that the (A) component and the (B) component dissolve mutually in order that using a solvent in process may bring about the conclusion of about [avoid] and crosslinking reaction, the structure of a bridge formation product, and the good repeatability of physical properties. (A) It is desirable to have the siloxane unit about the molecular structure and the chemical structure of the (B) component for dissolving with a component, and it is a general formula from the ease of formation of a macromolecule copolymer bridge formation object etc. [Formula 1]

$$R^{2}$$
 R^{2} R^{2} R^{2} R^{2} R^{2} R^{2} R^{2} R^{3} R^{2} R^{3} R^{2} R^{3} R^{3} R^{4} R^{3} R^{4} R^{3} R^{4} R^{4} R^{5} R^{4}

(-- an organic machine univalent in R2, an organic machine divalent in R3, an organic machine univalent [4/R] in an alkylene machine and R5, and a and c are / integer / of 2-1000 / and p of integer / of 0-1000 / and b / the integers of 2-100 among a formula The polyorganosiloxane which has polyoxyalkylene chain which has a hydroxyl group at the end shown by) as an at least 2 graft chain is desirable. As for this organopolysiloxane, aralkyl machines, such as aryl group; benzyls, such as alkyl group; phenyl groups, such as a methyl group, an ethyl group, and a propyl group, a tolyl group, and a xylyl group, and a phenethyl machine, are illustrated among an upper formula, as for R2. Moreover, a small amount of hydrogen atom and an alkoxy group may be contained in part. As for more than the moiety of R2, from a viewpoint of economical efficiency and the plasticity of a good macromolecule copolymer bridge formation object, it is desirable that it is a methyl group. R3 is divalent organic machines, such as arylene machines, such as an alkylene machine of the carbon numbers 1-8, such as a methylene group, an ethylene, a propylene machine, a butylene machine, a pentylene machine, a hexylene machine, a heptylene machine, and an octylene machine, or a phenylene group, and a naphthylene machine. R4 is alkylene machines, such as a methylene group, an ethylene, a propylene machine, a butylene machine, a pentylene machine, a hexylene machine, and a heptylene machine, R5 is acyl groups, such as an alkyl group; acetyl group or propionyl machines, such as a methyl group, an ethyl group, and a propyl group. Although a and c are within the limits of 0-1000, b is within the limits of 2-1000 and it is not limited especially, respectively, in order for the (A) component and the (B) component to dissolve, a and the ratio of (b+c) have desirable within the limits of - (1:5) (5:1). [0012] Although various methods are learned as the synthetic method of this graft copolymer The method of making carry out the graft of the polyoxyalkylene which has an unsaturation hydrocarbon group at the polyoxyalkylene which has an unsaturation hydrocarbon group at the piece end at the polyorganosiloxane which a part of side chain replaced by the hydrogen atom as the one method, and has an acyloxy machine at the other ends, and the piece end, and has a trimethylsilyl machine at the other ends by the hydrosilylation reaction with a predetermined ratio, and converting only the trimethylsilyl machine of a graft end into a hydroxyl group by the alcohol of an excessive amount after an appropriate time is mentioned.

[0013] (B) Although a component is a high molecular compound which has at least two hydroxyl groups in the above 1 molecules, and contains polyoxyalkylene chain in structure When higher ion conductivity is required This high molecular compound is general formula HO-(R6O) q-H (an alkylene machine and q of R6 are the integers of 1-100 among a formula.). Polyoxyalkylene or HO-(R7O) r-R8 (an organic machine univalent [7./ R] in an alkylene machine and R8 and r are the integers of 1-100 among a formula.) which has a hydroxyl group in the chain both ends shown It is desirable that it is polyoxyalkylene which has a hydroxyl group at the piece end of a chain shown. R6 and R7 of the upper formula of this polyoxyalkylene are alkylene machines, such as a methylene group, an ethylene, a propylene machine, a butylene machine, a pentylene machine, a hexylene machine, and a heptylene machine. R7 is acyl groups, such as an alkyl group; acetyl group or propionyl machines, such as a methyl group, an ethyl group, and a propyl group. q and r are within the limits of 1-100, and its within the limits of 5-20 is desirable. The condensation reaction of the polyoxyalkylene which has a hydroxyl group at the polyoxyalkylene or the piece end of a chain which has a hydroxyl group in these chain both ends is carried out to the above-mentioned (A) component,

and it carries out the work which increases the content of the bridge formation chain of polyoxyalkylene, or a graft chain into a macromolecule copolymer bridge formation object. Even if this bridge formation chain or a graft chain does not exist, the bridge formation chain or graft chain of polyoxyalkylene can be made to form by the polyoxyalkylene chain in a high molecular compound in this invention. Since there is a limitation in the amount of introduction of the polyoxyalkylene chain in a high molecular compound in the conditions which the (A) component and the (B) component dissolve, make it however, more desirable to contain the polyoxyalkylene which has a hydroxyl group at the polyoxyalkylene or the piece end of a chain which has a hydroxyl group in chain both ends in the (B) component. Moreover, in this invention, the direction where the graft chain of polyoxyalkylene existed a little in the macromolecule copolymer bridge formation object is in the inclination whose ion conductivity improves.

[0014] (C) Although a component is an alkali compound containing a lithium atom and especially the kind is not limited, a hydroxide, an alcoholate, a hydride, etc. are desirable and it is desirable to be used, especially the gestalt of a hydroxide, i.e., a lithium hydroxide, (LiOH). This (C) component carries out the role which finally forms the carboxyl group in the (A) component into lithium carboxylate by dehydration at the same time it carries out the catalysis of the esterification reaction of the (A) component and the (B) component. Consequently, as a macromolecule copolymer bridge formation object, it is fixed to a siloxane polymer chain and an anion (carboxylate ion) becomes the gestalt by which the cation (lithium ion) which is a counter ion was distributed. Moreover, as for the variance, it is desirable that ratio ratio [of the number of mols [Li+] of a lithium ion to the number of mols of the oxy-alkylene machine in a macromolecule copolymer bridge formation object [RO]] [Li+]/[RO] sets to 0.005-0.25, and it is 0.02-0.1 more preferably, this -- this ratio if 0.25 is exceeded --- the polarity of a macromolecule copolymer bridge formation object -- going up -- the maneuverability of a segment -- bad -- becoming -- moreover -- It is because ion conductivity high from the fall of the number of carriers will become difficult to get if it becomes less than 0.005. Moreover, the number of mols of the carboxyl group in the (A) component, the number of mols of the hydroxyl group in the (B) component, and the ratio of the sum total of the alkali equivalent in the (C) component are within the limits of - (1:10) (10:1), and its within the limits of - (1.0:1.2) (1.2:1.0) is desirable. Although the macromolecule copolymer bridge formation object with which the esterification reaction occurred and solidified (A) - (C) component by any ratio for this is obtained (B) When the ratios of the number of mols of the number of mols of the hydroxyl group in a component, the sum total of the alkali equivalent in the (C) component, and the carboxyl group in the (A) component differ sharply, be in the inclination which will be in the state where an unreacted carboxyl group or an unreacted hydroxyl group remained in solid electrolyte material. Since it is expected that evils, such as a reaction with electrode material, happen when the solid electrolyte in this state is included in a cell, it is not desirable. Moreover, if the above-mentioned mole ratio is set to 1, an anion kind will serve as only carboxylate ion to which only the lithium ion was fixed, and a cation kind will serve as a perfect lithium ion independent conduction-type solid electrolyte. [0015] Although the mixture which consists of a (A) component - (C) component in this method is stiffened by the dehydration condensation reaction and a macromolecule copolymer bridge formation object is made to form, this bridge formation object is mainly formed of the esterification reaction of the carboxyl group in the (A) component, and the hydroxyl group of the (B) component. Esterification reaction catalysts, such as a conventionally well-known reaction means, for example, alkali etc., by which it is used for the reaction of a carboxyl group and a hydroxyl group as the reaction means, are used, and the method of making it react to the bottom of ordinary temperature or heating can be used. Moreover, in order that heating may remove the water which is the by-product generated by the esterification reaction at the same time it promotes this esterification reaction, it is an effective technical means, and the temperature is usually 150 degrees C or less. Moreover, this crosslinking reaction can be performed on condition that a non-solvent. (A) A component and the (B) component are uniformly mixable easily by operation of stirring etc. (C) You may add, after making it dissolve in the (B) component beforehand or mixing the (A) component and the (B) component. since a component has solubility to an oxy-alkylene chain. (C) Although operation of stirring etc. is also possible for the dissolution to the oxy-alkylene chain of a component, for dissolution time

shortening, operation of heating or ultrasonic irradiation or addition of very little water is effective. Moreover, when use of the organic solvent is permissible in a dissolution process, a (A) component - (C) component may be mixed and dissolved in the organic solvent, and a solvent may be evaporated after an appropriate time. Although especially this organic solvent is not limited, a tetrahydrofuran, a dioxane, an acetonitrile, a JIMECHI formamide, and dimethyl sulfoxide are mentioned, for example. Moreover, since this esterification reaction generates water as a by-product, it is desirable to put the mixture of a (A) component - (C) component under reduced pressure finally, for example, the method of heating under reduced pressure, after advancing an esterification reaction to some extent under ordinary-pressure heating, and performing simultaneously the removal of water and the conclusion of an esterification reaction which were generated is recommended. However, when a solvent is used for a dissolution process, after advancing an esterification reaction to some extent below with the boiling point of the organic solvent under an ordinary pressure, the organic solvent is evaporated and it is necessary to heat under reduced pressure after an appropriate time.

[0016] Although the lithium cell of this invention comes to combine the above-mentioned positive electrode, a negative electrode, and a solid electrolyte, it is not limited especially about the building method. You may combine three persons, after producing each separately. However, since the efficiency as a cell increases so that the touch area in an interface is generally large, after carrying out the cast of the mixture of a solid electrolyte raw material on a positive electrode, making crosslinking reaction perform and film-izing, the method of making a negative electrode rival is recommended, in order that [moreover,] a negative electrode and a solid electrolyte may receive transformation with moisture -- production of a cell -- dry air -- it is necessary to carry out in the atmosphere of inert gas, such as an argon, preferably

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EXAMPLE

[Example] Hereafter, an example explains this invention to a detail more. In addition, measurement of the ionic conductivity of a solid electrolyte was performed by the following method. The solid electrolyte was cast in the shape of a film, and it considered as test sample. After measuring the thickness of this sample with a micrometer, the platinum electrode of the shape of a circular plate with a diameter of 1cm is stuck to both sides of a sample. After installing in the reduced pressure container which can set this whole as arbitrary temperature, decompressing to the high vacuum of 10 to 5 or less Torrs and the state of a sample fully reaching a balance, 5Hz - 13MHz alternating voltage was impressed by the LCR meter (4192made from YOKOGAWA ELECTRIC Hewlett Packard A), and conductivity was measured by the complex impedance method.

[Example 1] What is depended on the conventional technology (sample 2) was produced for the thing (sample 1) depended on this invention as a solid electrolyte, and property comparison. [0019] Stirring mixture of compound (1)0.384g, compound (2)0.450g, compound (3)0.167g, 12.2mg of lithium hydroxides, and 0.16g of water shown in the sample 1 following was carried out, the ultrasonic wave was irradiated, and it was made to fully dissolve. After slushing this solution into the pan made from the Teflon of 3cm around and heating it at 120 degrees C on a hot plate for 2 hours, when the vacuum drying was carried out for four days at 140 degrees C by the vacuum dryer, the film with the transparent thickness of 0.3mm was obtained. Since that the peak originating in a hydroxyl group and an isolation carboxylic acid is not accepted, the stretching-vibration peak of the carbonyl group by esterification being seen by 1740cm-1, and the unsymmetrical stretching-vibration peak by carboxylate ion were seen by 1600cm-1 when the infrared absorption spectrum of this film was investigated, it turns out that the esterification reaction is performed nearly completely. When the ion conductivity of a film was measured, the value of 2.0x10-7 S-cm -1 was acquired at 25 degrees C.

Compound (3): HO(CH2CH2O)12CH3 [0020] After having carried out stirring mixture of compound (1)0.278g, compound (2)0.527g, compound (3)0.195g, and 30.8mg of lithium perchlorates used by sample 2 sample 1, having irradiated the ultrasonic wave, making it fully dissolve and adding 3micro of decinormal hydrochloric-acid ethanol solutions l, when the same stoving processing as a sample 1

was performed, the film with the transparent thickness of 0.3mm was obtained. When the infrared absorption spectrum of this film was investigated, it turns out [that the peak originating in a hydroxyl group and an isolation carboxylic acid is not accepted, and / as which the stretching-vibration peak of the carbonyl group by esterification is regarded by 1740cm-1] that the shell and the esterification reaction are rattlingly performed nearly completely. When ion conductivity was measured, the value of 4.8x10-5 S-cm -1 was acquired at 25 degrees C.

[0021] The lithium foil with a diameter [of 1cm] and a thickness of 0.14mm was stuck to both sides of the sample 1 with a direct-current-voltage impression examination thickness of 0.3mm and a sample 2, this whole was installed in the reduced pressure container, after it decompressed to the high vacuum of 10 to 5 or less Torrs at 25 degrees C and the state of a sample fully reached the balance, the direct current voltage of 1V was impressed to both the lithiums foil, and flowing aging of current was observed. By the sample 1, 5.2micro of early current A continued for about two days, and it was confirmed that this ion conductivity material is cation independent conduction-type ion conductivity material. On the other hand, two days after by the sample 2, 79micro of early current A fell to 4.4microA. Subsequently, the lithium cell was constituted using the solid electrolyte of samples 1 or 2. The lithium-manganese multiple oxide used as a positive-electrode material was prepared by the method stated to Sanyo Technical Review and 20,114 (1989) by Noma and others. Lithium hydroxide After often mixing 0.118g and 1g of chemosynthesis manganese dioxide, it heated in 20-hour air at 375 degrees C, and the specified substance was obtained. When casting to an electrode, Teflon 5mg was added as acetylene black 20mg as 30mg of this lithium-manganese multiple oxide, and an electric conduction agent, and a binder, pressurization molding was carried out, and it obtained. The cross section of the cell produced using this positive electrode was shown in drawing 1. The chargeand-discharge cycle test was performed about the lithium cell with which samples 1 or 2 were incorporated. 3.5V and minimum voltage at the time of electric discharge were performed for the upper limit voltage at the time of charge as 2.5V by the constant current (3.77 or 37.7microA/cm -2). The voltage-time curve in a cycle typical to drawing 2 was shown. By the cell (a) incorporating the sample 1 obtained by this invention, not seeing and the inclination to go up further has most decline in the efficiency accompanying the repeat of charge and discharge. In the cell (b) incorporating the sample 2 obtained with the conventional technology, and the (example 1 of comparison), decline in the efficiency by the repeat of charge and discharge was seen. Transition of the service capacity for every cycle was shown in drawing 3. To having seldom changed the service capacity for every cycle by the cell (a) incorporating the sample 1, but being stable, in the cell (b) incorporating the sample 2, and the (example 1 of comparison), stability was missing and service capacity also fell gradually. [0022]

[Example 2] Compound (4)10g shown below was dissolved in about 500ml toluene / n-butyl alcohol (1/1) mixed solvent, the 76.42mg lithium hydroxide was added, it fully stirred, and the carboxyl group in a compound (4) was made to form into lithium carboxylate completely. Stirring mixture of compound (5)0.154g which removes a solvent completely after an appropriate time, and is shown in 0.772g of obtained oily matter and the following, and the compound (6)0.074g was carried out, the ultrasonic wave was irradiated, and it was made to fully dissolve. After adding about 20mg of phenyl-1-hide ROKISHI isopropyl ketones to this mixture as a photosensitizer (4-isopropyl), when it slushed into the pan made from the Teflon of 3cm around and the ultraviolet rays of 160 W/cm from a highpressure mercury lamp were irradiated for 6 seconds in 5cm distance from the perpendicular upper part, the film with the transparent thickness of 0.3mm was obtained. After carrying out reduced pressure drying of this for two days at 70 degrees C furthermore, when ionic conductivity was measured, the value of 1.6x10-7 S-cm -1 was acquired at 25 degrees C. Moreover, when the same direct-current-voltage impression examination as an example 1 was performed, the current of 4.7microA continued for about two days. When the still more nearly same lithium cell as an example 1 was produced, early service capacity is 2.5 mAh/g and became 20 cycle eye with 2.3 mAh/g. Compound (4): [Formula 4]

Compound (5): CH2=CH-CH2O(CH2CH2O)12CH2-CH=CH2 compound (6):CH2=CH-CH2O (CH2CH2O)12CH3 [0023]

[Example 3] Compound (7)10g shown below was dissolved in about 500ml toluene / n-butyl alcohol (1/1) mixed solvent, the 86.67mg lithium hydroxide was added, it fully stirred, and the carboxyl group in a compound (7) was made to form into lithium carboxylate completely. Stirring mixture of compound (5)0.170g which removed the solvent after an appropriate time completely, and used it for it in 0.749g of obtained oily matter and the example 1, and the compound (6)0.081g was carried out, the ultrasonic wave was irradiated, and it was made to fully dissolve. After 2.47microl Adding a chloroplatinic-acid 6 hydrate (H2PtCl6.6H2O) isopropyl alcohol solution to this mixture 2% of the weight as a hydrosilylation catalyst, it slushed into the pan made from the Teflon of 3cm around, and when heated within the oven adjusted to 70 degrees C for 2 hours, the film with the transparent thickness of 0.3mm was obtained. After carrying out reduced pressure drying of this for two days at 70 degrees C furthermore, when ionic conductivity was measured, the value of 1.8x10-7 S-cm -1 was acquired at 25 degrees C. Moreover, when the same direct-current-voltage impression examination as an example 1 was performed, the current of 4.9microA continued for about two days. When the still more nearly same lithium cell as an example 1 was produced, early service capacity is 3.5 mAh/g and became 20 cycle eye with 2.9 mAh/g.

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DESCRIPTION OF DRAWINGS

[Brief Description of the Drawings]

[Drawing 1] Drawing 1 is the cross section of the lithium cell of this invention.

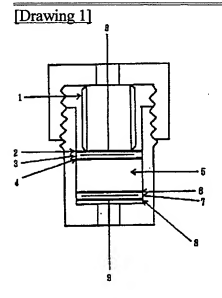
[Drawing 2] Drawing 2 a shows a time change of the voltage by the charge and discharge under the constant current (3.77microA/cm2) of the lithium cell obtained in the example 1, and drawing 2 b shows time change of the voltage by the charge and discharge under the constant current (37.7microA/cm2) of the example 1 of comparison. In addition, in drawing 2 a and drawing 2 b, v is voltage, and t is elapsed time.

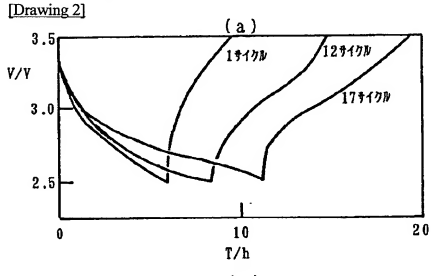
[Drawing 3] Drawing 3 a shows transition for every cycle of the service capacity by the charge and discharge under the constant current (3.77microA/cm2) of the lithium cell obtained in the example 1, and drawing 3 b shows transition for every cycle of the service capacity by the charge and discharge under the constant current (37.7microA/cm2) of the lithium cell of the example 1 of comparison. In addition, in drawing 3 a and drawing 3 b, D.C. is service capacity, and C.T. is the number of cycles. 1 -- the spacer made from Teflon, and 2 -- a nickel network and 3 -- a nickel network and 4 -- a positive electrode (lithium-manganese multiple oxide) and 5 -- a solid electrolyte and 6 -- a negative electrode (lithium foil) and 7 -- a nickel network (what stuck the lithium foil by pressure), and 8 -- a nickel network and 9 -- lead wire

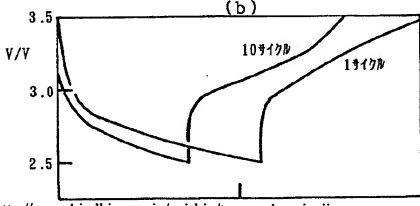
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DRAWINGS







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